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Luminescent Photoelectrochemical Cells. 7. Photoluminescent and Electroluminescent Properties of Cadmium Sulfo-Selenide Electrodes

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Samples of single-crystal, n-type  $CdS_XSe_{1-X}$  (X = 1.00, 0.74, 0.49, 0.11, 0.00) emit when excited with ultraband gap excitation. The 295 K band gaps monotonically decrease with X from  $\sim 2.4$  eV for CdS to  $\sim 1.7$  eV for CdSe. Photoluminescence (PL) spectra are sharp and have band positions which vary nearly linearly with composition:  $\lambda_{max}$  (nm) = 718-210 X. The energetic proximity to the band gap, temperature dependence and decay times are all consistent with a description of the PL as edge emission. Measured PL efficiencies,  $\phi_{\mathbf{r}}$ , are

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 $\sim 10^{-4}$  in air. When the samples are used as photoanodes in photoelectrochemical cells (PECs) employing aqueous polysulfide electrolyte, the emission intensity can be quenched by the passage of photocurrent. The extent of quenching can be correlated with the photocurrent quantum efficiency. Electroluminescence (EL) can be initiated by using the CdS $_{\rm X}$ Se $_{\rm 1-X}$  samples as dark cathodes in aqueous, alkaline, peroxydisulfate electrolyte at potentials cathodic of  $\sim -0.9$  to  $\sim -1.1$  V vs. SCE. The EL spectral distribution for a given sample is similar to that observed in PL experiments and indicates that the same emissive excited state is involved. At high resolution EL and PL spectra can differ in a manner which shows evidence of self-absorption effects: The EL spectra are slightly broader than their PL counterparts with the spectral mismatch almost exclusively in the high-energy tail. This suggests that EL occurs, on average, nearer the semiconductor-electrolyte interface. Measured EL efficiencies exceed  $10^{-4}$  (X = 0.00, 0.11) or  $10^{-5}$  (X = 0.49, 0.74, 1.00) at  $\sim -1.50$  V vs. SCE but are smaller near the EL threshold potentials. Comparisons with  $\sim 0.00$  values are

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Photoelectrochemical cells (PECs) are being widely studied as devices for optical energy conversion. (1). The excited-state properties of the semiconductors around which PECs are constructed are crucial to efficient energy conversion. We have employed luminescence as a probe of these excited-state properties, generally using materials such as n-type CdS:Te (Te-doped CdS) which exhibit subband gap emission (2).

Recently, we examined emission of band gap energy from n-type CdS and CdSe, two materials which have been used extensively in PEC studies. (1,3,4)

Since these two compounds form solid solutions over the entire composition range (5) the mixed compounds represent a natural extension of our emissive studies. We report herein that luminescence from samples of n-type, single-crystal  $CdS_{\overline{X}}Se_{\overline{1-X}}$  can be used to probe interfacial charge-transfer events relevant to PECs. Specifically, we demonstrate that photoluminescence (PL) can be perturbed and electroluminescence (EL) initiated by interfacial charge-transfer processes.

# Results and Discussion

Samples of n-type, single-crystal  $CdS_XSe_{1-X}$  representing several values of X were employed for the studies to be described, with CdS and CdSe included as reference points. In the sections which follow we characterize the composition of these samples and their PL and EL properties. PL characterization is based on the spectral distribution and origin of emission, radiative efficiency and PEC properties. EL characterization includes spectral distribution and efficiency.

Sample Composition. Two methods which have previously been used to characterize the composition of cadmium sulfo-selenide samples are electron microprobe analysis (6) and absorption spectra (7). For the five  $CdS_XSe_{1-X}$  samples employed

in this study, we found by using an electron microprobe analyzer that X was 1.00, 0.74, 0.49, 0.11, and 0.00; we estimate these compositions to be accurate to better than 1%. Corresponding to these values of X are crystal colors of yellow, red, dark red, black, and black. The trend in color reflects direct band gaps which are reported to monotonically decrease with increasing Se content (6-8). Figure 1 displays the red shift in absorption edge with increasing Se concentration for the five  $CdS_XSe_{1-X}$  samples investigated in this study. Good agreement is observed with spectra previously reported for several of these samples (7b,9,10). Crude estimates of band gap energies,  $E_{BC}$ , from the data of Figure 1 are  $\sim 2.37$ , 2.10, 1.93, 1.72 and 1.68 eV for X = 1.00, 0.74, 0.49, 0.11 and 0.00, respectively, using an absorptivity,  $\alpha$ , of 10 cm<sup>-1</sup> as a comparison point. These values are in reasonable accord with the various estimates reported from measurements of absorption spectra (7), photoconductivity spectra (8a,b), and PEC photoaction spectra (11); plots of  $E_{BC}$  (eV) vs. X are in all cases sublinear with respect to a line drawn between the CdS and CdSe  $E_{RC}$  end points.

PL Spectra and Origin. When the CdS XSe<sub>1-X</sub> samples are irradiated with ultraband gap light, they visibly luminesce. Corresponding to the X values of 1.00, 0.74, 0.49, 0.11 and 0.00, the emitted light is green, yellow-green, orange, red and red. Uncorrected front-surface emission spectra of the samples were obtained in 5M OH with 457.9-nm excitation while they served as electrodes, Figures 2a and 3a, and are quite similar to spectra obtained in air. The bands in these figures are all fairly sharp (fwhm values of ~0.05-0.07 eV) and their position red shifts with increasing Se content in accord with the observed colors. A plot of the emission band maxima (nm) vs. X is presented in Figure 4. The observed roughly linear relationship given by eq. (1) is similar to that previously reported in cathodoluminescence and

$$\lambda_{\text{max}}(\text{nm}) \approx 718-210 \text{ X}$$
 (1)  
for  $\text{CdS}_{X}\text{Se}_{1-X} (0 \le X \le 1)$ 

photoluminescence studies (6). As noted in the earlier study, eq. (1) permits an independent determination of composition in  $CdS_XSe_{1-X}$  samples.

Comparison of Figure 1 with Figures 2a and 3a indicates that the emission bands of the five samples overlap the onset of their absorption. The overlap is occasionally also reflected in the shapes of the emission bands: High resolution spectra of several  $CdS_XSe_{1-X}$  samples exhibit breadths which can be dependent on excitation wavelength because of self-absorption effects. For example, the PL spectrum of CdSe from 457.9 nm excitation is broader than that obtained from  $\sim 3$ -fold more deeply penetrating 632.8 nm excitation ( $\sim 22$ -vs. 19-nm fwhm values) with the spectral mismatch occurring almost exclusively in the high-energy tail of the emission band where the probability for reabsorption of the emitted light is greatest (4); more modest but analogous differences are found in comparing PL spectra obtained with 457.9 and 514.5 nm excitation for the X = 0.00, 0.11, 0.49, and 0.74 samples. Self-absorption effects are also observed in comparisons of PL and EL spectra (vide infra).

The energetic proximity of the emission bands to the band gap region suggests that the PL be described as edge emission (12). Although we are uncertain as to the exact origin of these radiative transitions in our samples, temperature effects and decay times support this description. Cooling the samples to 77 K results in a marked blue shift of the emission band maxima. The magnitude of these shifts is given in Table I and generally ranges from  $\sim 0.07-0.10$  eV. This energetic difference roughly matches the increase in  $E_{BG}$  reported for  $CdS_XSe_{1-X}$  samples from photoconductivity measurements (8) and supports the notion that the transition is related to  $E_{BG}$ . At 77 K we found that several of the samples also exhibited subband gap emission bands. These bands are listed in Table I, but we have not characterized them further. Decay times of all five samples were measured at 295 K in air using 337 nm excitation from a  $N_2$  laser; incident intensities of  $\sim 3$  and 30 kW/cm<sup>2</sup> were used and gave similar results. In accord with the literature (13), a decay time of  $\sim 20$  ns

 $(\tau_{1/e})$  was observed for CdSe. The decay time of the X = 0.11 sample was also  $\sim$ 20 ns, but those of the other samples were limited by the laser pulse duration and are  $\lesssim$  7 ns. Although our inability to obtain more precise values is disappointing, decay times in the (sub)nanosecond regime have been reported for edge emission (14). Taken together, we feel that the energies, temperature dependence and decay times of the observed PL bands are consistent with their classification as edge emission.

PL Efficiency. The PL quantum efficiency, defined as photons emitted per photons absorbed and symbolized by  $\phi_r$ , is difficult to measure accurately. An upper limit to  $\phi_r$  can be obtained for a given set of experimental conditions by finding other conditions which increase its value. As Table I indicates, cooling the CdS $_X$ Se $_{1-X}$  samples to 77 K not only blue shifts the highest energy emission band but also markedly increases its intensity. Emission enhancement factors of  $\sim$ 20 to 200 correspond to 295 K upper limits on  $\phi_r$  of 0.05 to 0.005; we assume little change in absorbed 457.9-nm intensity in this experiment.

A second estimate of  $\phi_r$  is based on a technique developed by Wrighton, et al (15). The "photons absorbed" needed for the  $\phi_r$  calculation is acquired by subtracting the intensity of laser light reflected by the  $CdS_XSe_{1-X}$  sample from the intensity reflected by a nonabsorbing standard, a MgO pellet, in the same geometry. This difference is divided into the intensity emitted by the  $CdS_XSe_{1-X}$  sample to obtain  $\phi_r$  (see Experimental). Typical  $\phi_r$  values are  $\sim 10^{-4}$ , consistent with the aforementioned upper-limit estimate. We have occasionally seen  $\phi_r$  as large as 4 x  $10^{-4}$  for CdSe by this technique which is estimated accurate to  $\pm 25\%$  (15). Our absolute values will be scmewhat low, however, owing to self-absorption.

PEC Properties. Although inefficient, luminescence provides insight into the effects of PEC parameters on the excited-state properties of semiconductor electrodes. We constructed one-compartment PECs in the sample compartment of

an emission spectrometer. Besides the  $CdS_XSe_{1-X}$  photoanode, the PEC consisted of a Pt foil counterelectrode, an SCE, and an aqueous polysulfide electrolyte whose composition was 1 M  $\mathrm{OH}^-/\mathrm{1M}\,\mathrm{S}^{2-}/\mathrm{0.1}\,\mathrm{M}\,\mathrm{S}$ . Examination of the front-surface luminescence in these PECs was facilitated by the insensitivity to potential of the emissive spectral distribution (2.0-nm resolution); between the onset of photocurrent and -0.3 V vs. SCE essentially only the emission intensity changed, permitting it to be monitored by simply sitting at  $\lambda_{max}$ . PL spectra in polysulfide electrolyte were virtually identical to those obtained in OH electrolyte shown in Figures 2a and 3a. The interrelationship of photocurrent, luminescence, and voltage for  $CdS_{\chi}Se_{1-\chi}$ -based PECs is presented in the iLV curves of Figure 5 which were all obtained with 457.9-nm light. At this excitation wavelength the photogenerated electron-hole (e - h +) pairs are formed almost exclusively in the depletion region where they are most susceptible to potential-induced changes in band bending (16). Figure 5 reveals that all five semiconductor electrodes exhibit decreasing photocurrents and increasing emission intensities with increasingly negative potentials. This inverse relationship is anticipated for what are essentially competitive excited-state deactivation processes: photocurrent is a measure of e - h pair separation, whereas luminescence is a probe of their recombination. Increasingly negative bias reduces band bending for n-type semiconductors (16); the decreased electric field inhibits e - h pair separation and promotes their recombination, as illustrated in Figure 5.

A relationship between photocurrent and emission intensity examined through iLV curves of CdS:Te-based PECs is given by eq. (2) (17,18). The symbols  $\phi_{\bf x}$ ,  $\phi_{\bf r}$ ,

$$\frac{\phi_{r_o}}{\phi_{r_i}} = \frac{1}{1 - \phi_{x}} \tag{2}$$

and  $\phi$  represent quantum efficiencies for photocurrent, open-circuit and in-circuit i radiative recombination, respectively. The key assumption made in deriving eq. (2) is

that the ratio of  $e^- - h^+$  pairs recombining radiatively to those recombining nonradiatively is unaffected by potential. The iLV curves of Figure 5, like those of CdS:Te-based PECs, are in reasonable accord with eq. (2). For example, the emission for the X = 0.74 electrode yields a  $\phi_r$  / $\phi_r$  ratio of  $\sim$ 15, using -0.3 V vs. SCE as the  $\phi_r$  point. From eq. (2)  $\phi_x$  is predicted to be 0.93, in good agreement with the direct measurement of 0.91; the direct measurement is uncorrected for reflection and electrolyte absorption. Similar comparisons for the other iLV curves of Figure 5 are given in Table II and support the applicability of eq. (2) to CdS $_x$ Se $_{1-x}$ -based PECs.

Also presented in Table II are energy conversion characteristics of the Figure 5 curves. Monochromatic optical energy conversion efficiencies of ~7-10% were obtained for the single-crystal mixed samples and are comparable to values reported for polycrystalline samples used in similar PECs (11). We should mention, too, that although no changes in the iLV curves were observed after several scans, exchange reactions between the electrode and electrolyte (poly)chalcogenide species have been reported and would be expected to alter the curves over longer experimental periods (19).

EL Mechanism and Spectra. EL was studied in a  $N_2$ -purged, one-compartment cell consisting of a  $CdS_XSe_{1-X}$  working electrode, a Pt counterelectrode, an SCE reference electrode and an aqueous, alkaline peroxydisulfate electrolyte (5 M  $OH^-/0.1$  M  $S_2O_8^{-2-}$ ). When sufficiently cathodic potentials were applied to any of the  $CdS_XSe_{1-X}$  electrodes, luminescence of the color emitted by that electrode

in PL experiments was observed. The mechanism proposed for EL is given in eq. (3)-(5) (20):

$$s_2 o_8^{2-} + e_{CB}^{-} \longrightarrow so_4^{2-} + so_4^{-}$$
 (3)

$$so_4$$
:  $\longrightarrow so_4^{2-} + h^+_{VB}$  (4)

$$e^{-} + h^{+} \longrightarrow 1ight$$
 (5)

In this scheme an electron in or near the conduction band  $(e_{CB}^-)$  reduces  $s_2 o_8^{-2}$  to yield the strongly oxidizing  $so_4^-$  radical; the radical then injects a hole into or near the valence band  $(h_{VB}^+)$ . Subsequent radiative recombination yields emission of band gap or subband gap energy. The former case obtains if both the electron and hole are near the band edges; if either or both species are located in intraband gap states such as surface states or dopant states, subband gap emission results.

The redox potentials of eq. (3) and (4) have been estimated to be  $\leq +0.6$  and  $\geq +3.4$  V vs. NHE, respectively (19b). Initiation of the reaction sequence of eq. (3)-(5) should occur at potentials cathodic of the flat-band potential,  $E_{FB}$ , where the  $CdS_XSe_{1-X}$  samples can serve as dark cathodes. We estimate that  $E_{FB}$  values for the samples in aqueous alkaline solution are in the range of  $\sim -0.7$  to -1.0 V vs. SCE. This range is based on values reported for CdS and CdSe and i-V properties of polycrystalline  $CdS_XSe_{1-X}$  samples (11, 21). Figure 6 illustrates the i-V properties of the mixed sulfo-selenide electrodes in CH and OH  $S_2O_8^{2-}$  electrolytes. Consistent with the postulated  $E_{FB}$  position, cathodic current onsets between  $\sim -0.8$  and -1.0 V vs SCE. For potentials  $\geq -1.6$  V the current is due predominantly to reduction of  $S_2O_8^{2-}$ . Similar i-V properties were observed for CdS and CdSe electrodes (4, 22).

Another electrochemical process which can contribute to the observed current is reduction of the electrode to yield surface Cd, eq. (6). This reaction has been reported for CdS in neutral media and

$$CdS_XSe_{1-X} + 2e^- \longrightarrow Cd + XS^{2-} + (1-X)Se^{2-}$$
 (6)

can be photoassisted (23a); evidence for the reaction was the appearance, after reduction, of a sharp anodic stripping peak accompanied by changes in electrode reflectivity (23a) and photoacoustic signal (23b). In OH or OH  $/S_2O_8^{2}$  electrolytes, we generally did not observe an anodic stripping peak in dark i-V curves (0.0 to -1.5 V) of  $CdS_XSe_{1-X}$  samples. That a surface layer might be present, however, was suggested by decays in current and EL intensity which were observed when the  $CdS_XSe_{1-X}$  electrodes were brought into circuit in OH or OH  $/S_2^{0}$  electrolytes at potentials cathodic of  $\sim$ -1.2 V. If electrode reduction were occurring, earlier studies on CdS suggested that anodic cycling might reverse such surface changes (20c,22,23). Evidence that reproducible i-V properties could be obtained in  $0 \text{ h}/\text{S}_2 \text{O}_8^{2}$  electrolyte was indicated by pulse experiments: repetitively pulsing the  $CdS_{\chi}Se_{1-\chi}$  electrodes between 0.0 V (1s) and -1.5 V (1s) gave reproducible coulombs during the cathodic cycle over hundreds of pulses. Additionally, as with CdS (22) and CdSe (4), little weight loss was found in long-term pulse experiments: For example, a 0.1275 g  $^{\text{CdS}}_{0.49}^{\text{Se}}_{0.51}$ sample ( $\sim$ 0.4 cm $^2$  surface area) showed negligible weight loss after 145 C had been passed in ∿14 hrs of continuous pulsing between 0.00 V (1s) and -1.50 V (1s) in 10 ml of OH /S202 electrolyte. Although these results could be ascribed to reversal of electrode reduction during the anodic cycles, any such conclusions await a detailed characterization of the electrochemistry of  $CdS_XSe_{1-X}$  electrodes in  $OH^{-}/S_{2}O_{8}^{-2-}$  electrolyte. The possible presence of a surface layer should be kept in mind, however, since it could influence the EL properties to be described.

Placement of the EL cell in the sample compartment of the emission spectrometer permitted EL spectra to be recorded. Threshold potentials for visibly observing EL from the  $CdS_XSe_{1-X}$  electrodes range from  $\sim -0.9$  to -1.1 V vs. SCE. Because EL intensities generally decayed from their initial value, the pulse technique was employed to obtain EL spectra:  $CdS_XSe_{1-X}$  electrodes were repetitively pulsed between 0.00 V (11s) and a potential cathodic of the EL threshold potential (1s) while the emission monochromator was slowly scanned. The transient maximum intensities obtained in this procedure were reproducible ( $\pm 10\%$ ) over hundreds of pulses. EL spectra were typically recorded under conditions which facilitated comparisons with PL spectra. For a given sample the PL spectrum was first obtained in air and at -1.50 V in  $OH^-$  electrolyte. Without disturbing the sample geometry, the electrolyte was changed to  $OH^-/S_2O_8^{2-}$  and the EL spectrum obtained by pulsing between 0.00 and -1.50 V vs. SCE. These pairs of EL and PL spectra are given in Figures 2 and 3. The similarity of the PL and EL spectra indicate that a common emissive excited state is involved in both experiments.

Spectral comparisons made at higher resolution (% 0.3 nm as opposed to 2.0 nm in Figures 2 and 3) often revealed spectral mismatches, as illustrated in Figure 7. The EL spectrum of  $\text{CdS}_{0.49} \text{Se}_{0.51}$  is slightly broader than the PL spectrum (fwhm of 22 vs. 20 nm) with the discrepancy occurring almost exclusively in the high-energy tail. We attribute the difference in EL and PL spectra to self-absorption effects. The discrepancy in the breadths of the two spectra of Figure 7 indicates that, on average, EL is produced nearer to the  $\text{CdS}_{0.49} \text{Se}_{0.51}$  electrolyte interface than PL. Similar effects were observed for the other  $\text{CdS}_{X} \text{Se}_{1-X}$  samples with the magnitude of the spectral discrepancy dependent on sample and PL excitation wavelength. Quantifying these inferred differences in EL and PL spatial origins should be possible, in principle, but requires more information than we presently possess. Among the data needed are the spatial distribution of injected holes, their diffusion length, and evaluation of the role, if any, or a surface layer (vide supra). For the mixed sulfo-selenide samples high-resolution absorption spectra are also

A final point regarding the EL spectra involves subband gap emission. Broad emission bands below the band gap energy have been observed in EL spectra of CdS (20c,22). At the sensitivities used in Figures 2 and 3, the EL spectra of the X = 1.00 and 0.74 samples were characterized by barely discernible emission between ∿600-800 nm in addition to edge emission. We did not observe subband gap emission from the other CdS Se samples at this sensitivity for  $\lambda < 800$  nm, the instrumental wavelength limitation. As described above, the presence of subband gap emission implicates the involvement of intraband gap states in the radiative transition. Although these states are presently uncharacterized for the systems at hand, our inability to detect the corresponding emission bands in PL experiments conducted under similar conditions (5M OH electrolyte; -1.50 V vs. SCE) is noteworthy. We found that homogeneously doped samples of CdS: Te exhibited subband gap emission in both PL and EL spectra (22). The observed difference in PL and EL spectra for the X = 1.00 and 0.74 samples may reflect an inhomogeneous distribution of the intraband gap states. In particular, localization of the intraband gap states at or near the surface (surface states, e.g.) would make detection of the corresponding emission easier in an EL experiment, if EL were significantly more surface-sensitive than PL. Although our data suggests that this might be the case, additional experiments are needed to prove it conclusively. EL Efficiency. We have adapted measures of efficiency from studies of electrogenerated chemiluminescence (ecl) to make estimates of EL efficiency (24). Integrated EL efficiencies,  $\overline{\phi}_{\text{EL}}$ , were determined by the pulse technique described earlier in conjunction with eq. (7); F is Faraday's constant. Total emitted light was estimated

$$\overline{\phi}_{EL} = \frac{\text{total emitted light (einsteins) x F}}{\text{total coulombs due to hole injection}}$$
(7)

by placing a radiometer next to the sole exposed face of the emitting electrode.

The integrated energy in a single-pulse experiment ( $\mu$ J) was converted to einsteins at  $\lambda_{max}$  of the emission band for each of the  $CdS_XSe_{1-X}$  samples; the sharpness of the bands makes this a reasonable approximation. Since not all of the photons are collected in this procedure, the value used for the numerator of eq. (7) is a lower limit. The denominator of eq. (7) was obtained by using a digital coulometer to integrate the coulombs over the single pulse. Because we use total coulombs for the calculation and not the coulombs due to hole injection, we are likely overestimating the denominator of eq. (7). If equations (3)-(5) are mechanistically correct, for example, our use of total coulombs is too large by a factor of two. Additionally, we have not corrected for background coulombs obtained in the absence of  $S_2O_8^{2-}$ , although this is at most a few percent for potentials  $\geq$ -1.5 V vs. SCE. The combination of underestimated numerator and overestimated denominator in eq. (7) yields  $\overline{\phi}_{EL}$  estimates which should be treated as lower-limit estimates.

Table III presents a compilation of  $\overline{\phi}_{EL}$  values near the EL threshold potential and at -1.50 V vs. SCE for the CdS $_X$ Se $_{1-X}$  samples. At -1.50 V the EL efficiencies range from  $\sim 10^{-5}$  to in excess of  $10^{-4}$  with the two samples having the highest Se content consistently giving the largest  $\overline{\phi}_{EL}$  values. Only modest declines in  $\overline{\phi}_{EL}$  (factors of  $\sim 2$ -8) are observed in passing to the threshold potential, although the constituents of  $\overline{\phi}_{EL}$  ( $\mu J$ , coulombs) often decreased by much larger factors.

Comparisons of  $\phi_r$  and  $\overline{\phi}_{EL}$  are noteworthy but must be made with care. The value of  $\phi_r$  measured in air for the  $\operatorname{CdS}_X \operatorname{Se}_{1-X}$  samples,  ${\sim}10^{-4}$ , is comparable to the  $\overline{\phi}_{EL}$  values of Table III. A more meaningful comparison can be made, however, if  $\phi_r$  is determined in OH electrolyte at the same potentials used in EL experiments. We found for CdSe, in fact, that lower-limit estimates of  $\phi_r$  were larger,  ${\sim}1 \times 10^{-3}$  at  ${\sim}1.50$  V in 5 M OH, but that this value dropped by a factor of  ${\sim}3$  by  ${\sim}0.9$  V (4). Figure 8 presents a plot of relative emission intensity vs. Fotential for several of the  $\operatorname{CdS}_X \operatorname{Se}_{1-X}$  samples including CdSe. If there is little change

in absorbed intensity over the -1.50 to -0.90 V potential excursion, then Figure 8 also represents relative  $\boldsymbol{\varphi}_{_{\boldsymbol{r}}}$  values as a function of potential. The decline with anodic potential is similar to the trend in measured  $\overline{\phi}_{FI}$  values with potential. This relationship is intriguing because, by analogy with ecl studies (24),  $\overline{\phi}_{EL}$  can be factored into an efficiency for excited-state population,  $\phi_{\text{FS}}$  (excited states populated per holes injected), and the radiative efficiency,  $\phi_r$  (photons emitted per excited states populated). Although the uncertainties in  $\boldsymbol{\varphi}_{\mathbf{r}}$  and  $\overline{\boldsymbol{\varphi}}_{EL}$  preclude a more precise comparison, it is tempting to conclude that  $\varphi_{\mathbf{F},\mathbf{C}}$  is very large, perhaps approaching its maximum value of unity, over the potential range examined. However, the different spatial origins for PL and EL inferred from their spectra suggest caution in comparing efficiencies, even when  $\phi_{r}$  and  $\overline{\phi}_{rl}$  are known with certainty. Strictly speaking, PL and EL efficiencies should only be compared in the same spatial regions of the electrode, since  $\phi_{_{\mathbf{r}}}$  values may be region dependent because of local variations in environment. The solid-liquid interface is a particularly important environment for EL, since it is the region in which EL is initiated and EL appears to occur, on average, nearer to it than PL. We have, in fact, seen substantial discrepancies in our estimates of  $\overline{\phi}_{EL}$  and  $\phi_r$  near the EL threshold potential for CdSe and this could be attributed to changes in  $\phi_{ES}$  and/or  $\phi_{r}$  resulting from altered (near-)surface conditions (4). To summarize this section, until the spatial dependence of  $\phi_{rs}$  and  $\phi_{r}$  can be determined more accurately, only limited comparisons between EL and PL efficiencies can be made.

In conclusion, we feel that this study demonstrates several useful applications of luminescence to the characterization of  $CdS_XSe_{1-X}$  electrodes. Perturbation of PL and initiation of EL through interfacial charge-transfer processes is shown to be a general feature of these materials and provides insight into the population and deactivation of the electrodes' emissive excited states. Comparisons

of PL and EL spectral distributions and efficiencies highlight spatial and mechanistic features of the two kinds of emission. Experiments to better understand the interrelationship of PL, EL and charge-transfer events are in progress.

## **Experimental**

Materials. Single-crystal c-plates (10x10x1 mm) of n-type  $CdS_XSe_{1-X}$  (X = 1.00, 0.74, 0.49, 0.11, and 0.00) were obtained from Cleveland Crystals, Cleveland, Ohio. The crystals were vapor-grown with resistivities of ~2 ohm-cm. After being cut to dimensions of  $\sim 0.25 \text{ cm}^2 \times 1 \text{ mm}$ , the samples were etched in Br<sub>2</sub>/MeOH (1:10 v/v) and mounted as electrodes as described previously (2b). The preparation of polysulfide and peroxydisulfate electrolytes has also been described (2b 22). MgO powder was obtained from Baker Chemical Co., MnBr<sub>2</sub>·4H<sub>2</sub>O from Aldrich, and Et<sub>4</sub>NBr from Eastman. Optical Measurements. Absorption spectra were obtained on a Cary 17-D spectrophotometer. Uncorrected emission spectra were taken with an Aminco-Bowman spectrophotofluorometer (200-800 nm; band width ∿1.0 or 2.0 nm) equipped with a Hamamatsu R446S red-sensitive PMT; spectra were displayed on a HP7004A x-y recorder. A McPherson Model 270, 0.35-m monochromator, equipped with both a grating blazed at 500 nm and the R446S PMT, was employed for high-resolution (~0.3-nm bandwidth) spectra. A grating blazed at 1000 nm and an EMI Model 9684B PMT (S-1 response) extended the spectral range to 1100 nm. Signals from either PMT used in the McPherson-based spectrometer were amplified by a Keithley Model 414S picoammeter and displayed on a Houston Model 2000 x-y recorder. The 457.9- and 514.5-nm lines of a Coherent Radiation CR-12 Ar ion laser were used for excitation. The 2-3-mm diameter beam was 10X expanded and passed through either a Melles Griot 03-FCG055 long-pass filter (cutoff ∿380 nm) or interference filter for 457.9- or 514.5-nm excitation, respectively, to eliminate laser plasma lines. The beam was then translated by periscope into the sample compartment of the Aminco-Bowman spectrometer; no periscope was needed when using the McPherson-based spectrometer.

The laser beam was often masked to fill the sample surface area. Laser excitation was filtered in emission spectral measurements by placing a Corning 3-66 filter or ferricyanide filter solution (0.03 M K<sub>3</sub>Fe(CN)<sub>6</sub>/0.6 M HCl; 1.0-cm pathlength cell) for 514.5- or 457.9-nm excitation, respectively, in front of the PMT. Laser intensity was adjusted by changing the power or by neutral density filters. The intensity was measured with a Scientech 362 power energy meter (flat response, 250-35,000 nm) or a Tektronix J16 radiometer equipped with a J6502 probe head (flat response, 450-950 nm). An EG & G Model 550-1 radiometer (flat response, 460-975 nm) equipped with a Model 550-3 pulse integration module was used for integrated energy measurements of the light produced in pulse EL experiments. A quartz disk was used as a beam splitter during experiments requiring continuous monitoring of laser intensity.

Sample Composition. Absorption spectra were obtained from samples which had been polished sequentially with 5.0-and 1.0-µm alumina. Samples were placed in a metal holder which restricted the optical window to a  $^{\circ}$ 2 mm diameter hole. X-ray analysis was performed on a Bausch & Lomb ARL SEMQ electron microprobe. Samples were ohmically contacted with Ga/In eutectic and Ag epoxy to Al discs. The crystals were mechanically fastened with epoxy and polished with 1.0 µm alumina to obtain flat samples. The primary electron beam energy of 15 keV yielded a sample current of 20 nA. All concentrations were corrected by a standard ZAF correction program. At least 5 regions of each sample were examined to establish sample homogeneity.

PL Properties. Front-surface PL spectra were generally obtained by positioning the sample at ~45° to both the Ar ion laser beam and the emission detection optics. The spectral mismatch described in the text was observed for several sample geometries including "head on" excitation along the c-axis (vide infra) and, for some samples, also using a CR Model 80 He-Ne laser for 632.8 nm excitation. Spectra were taken both in air and with the samples serving as electrodes in 5 M OH at various potentials. Spectra at 77 K were obtained

by placing samples in a liquid N<sub>2</sub>-cooled quartz Dewar which was positioned in the sample compartment of the Aminco-Bowman spectrometer as described previously (2b); spectra beyond 800 nm were taken on the McPherson-based spectrometer. Relative emission intensities between 295 and 77 K which were needed for upper-limit radiative efficiencies required corrected spectra. We determined crude correction factors for the Aminco-Bowman spectrometer as follows: We first measured the output of the high-pressure Xe lamp which is part of the spectrometer with the Model 550-1 radiometer; the output was measured from 400-800 nm (2.0-nm bandwidth) using the spectrometer's excitation monochromator and expressed in relative einsteins. Subsequently, the monochromatized output of the Xe lamp was reflected off a mirror into the emission detection optics and the detected intensity recorded at 10-nm intervals. The correction factors thus acquired were applied to areas under the edge emission bands at 77 and 295 K to yield an upper limit to  $\phi_{_{\mbox{\scriptsize T}}}$  at 295 K. A second estimate of  $\phi_r$  was made by placing the  $CdS_X Se_{1-X}$  samples in a quartz cuvette positioned in the Aminco-Bowman spectrometer's sample compartment as previously described (15). The central portion of the sample was irradiated "head-on" with the masked, expanded laser beam. Some of the emitted and reflected incident light was deflected into the detection optics by a mirror. The emission spectrum was then recorded with a calibrated neutral density filter present while scanning the excitation region; the filter was removed while scanning the emission band. At this point the sample was replaced by a highly reflective MgO pellet (pressed at 12,000 lbs.) and the excitation line recorded using an appropriate neutral density filter. The difference in areas of the reflected light between MgO and the  $\operatorname{CdS}_{\chi}\operatorname{Se}_{1-\chi}$  sample represents photons absorbed by the semiconductor and is divided into the area under the emission band, representing photons emitted. Both the absorbed and emitted intensities are corrected for relative detector response.

The salt  $[{\rm Et}_4{\rm N}]_2[{\rm MnBr}_4]$ , prepared as described in the literature (25), served as a standard and gave a  $\phi_r$  value similar to that previously reported (15). Decay times for  ${\rm CdS}_X{\rm Se}_{1-X}$  samples were obtained in air with 337-nm excitation using instrumentation and techniques previously reported (3). Peak intensities of  $\sim 3$  and 30 kW/cm<sup>2</sup> were used with essentially no difference in the resulting decay curves.

PEC Experiments. A cell suitable for conducting PEC studies in polysulfide electrolyte has been described (2b). Potentiostatic iLV curves were obtained using a PAR Model 173 potentiostat/galvanostat, a Model 175 programmer, and a Model 179 digital coulometer/I to E converter; the techniques used to generate these curves and to measure photocurrent quantum efficiency have been reported (2b). The Aminco-Bowman spectrometer (2.0-nm resolution) was used exclusively for obtaining iLV data with the emission intensity monitored at the emission band maxima of the CdS<sub>X</sub>Se<sub>1-X</sub> samples.

EL Current-Voltage Properties. Current-voltage curves relevant to EL studies were taken in both 5 M OH and 5 M OH /0.1 M S $_2^{0}$ 0 $_8^{2}$ . The curves were swept in a cathodic direction at 20 mV/s, beginning at 0.00 V vs. SCE. Electrolytes were stirred by a N $_2$ -purge.

EL Spectra. Pulse EL spectra were obtained in both the Aminco-Bowman and McPherson-based spectrometers; experiments in the Aminco-Bowman instrument have been described previously (22). Pulsing the electrode between 0.00 V (11s) and potentials cathodic of ~ -1.0 V (1s) gave sufficiently intense and reproducible transient signals (recorder response time of 0.3 s) to permit the EL spectrum to be obtained as a series of vertical lines as the emission monochromator was swept. Direct comparisons of EL and PL spectra were made for the CdS<sub>X</sub>Se<sub>1-X</sub> compounds: For each sample the low-resolution PL spectrum (2.0-nm bandwidth; Aminco-Bowman) was initially obtained in air and in 5 M OH electrolyte at -1.50 V vs. SCE using 457.9-nm excitation. Without disturbing the sample geometry, the electrolyte was

changed to 5 M OH  $\frac{1}{0.1}$  M S<sub>2</sub>0<sub>8</sub> and the EL spectrum taken by pulsing between 0.00 and -1.50 V, as described above. For high-resolution spectra (0.3-nm bandwidth; McPherson) essentially the same procedure was used, except that the sequence was PL in air, EL, and PL in the OH electrolyte; no change in sample geometry occurred throughout the sequence. For some samples 514.5- rather than 457.9-nm light was used for the PL spectra. It was possible to obtain for CdSe a steady-state EL spectrum at -1.30 to -1.50 V at high resolution and no difference with the pulse-generated EL spectrum was observed for the same sample geometry. EL Efficiency. The integrated EL efficiency,  $\overline{\phi}_{EL}$ , was estimated by placing the probe head of the EG & G radiometer as close as possible (within ∿l cm) to the sole exposed  $CdS_XSe_{1-X}$  crystal face. The electrode was pulsed to a given potential for 1s as described above and the total emitted energy (in µJ) per pulse measured with the EG & G Model 550-3 accessory. Total coulombs passed during the pulse were measured with the digital coulometer. These measurements were repeated as a function of potential. It should be mentioned that the position of the radiometer required the removal of its flat-response window. Consequently, measured values were corrected by factors determined at larger distances where the window could be added and removed without disturbing the relative sample-radiometer geometry; the correction factors used for the  $CdS_{\gamma}Se_{1-\gamma}$  samples were in good agreement with the manufacturer's values for the various wavelength regions involved. Finally, relative PL intensity as a function of potential (457.9-nm excitation) was obtained in 5 M OH electrolyte for comparison with  $\overline{\phi}_{EL}$ . PL intensity was monitored at  $\lambda_{max}$  of the CdS $_X$ Se $_{1-X}$ sample while the electrode potential was swept between -0.90 and -1.50 V vs. SCE at 20 mV/s.

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#### References

- (a) A. J. Bard, <u>Science</u>, <u>207</u>, 139 (1980); (b) M. S. Wrighton, <u>Acc. Chem.</u>
   Res., <u>12</u>, 303 (1979); (c) A. J. Nozik, <u>Ann. Rev. Phys. Chem. <u>29</u>, 189 (1978).
  </u>
- (a) A. B. Ellis and B. R. Karas, <u>Adv. Chem. Ser.</u>, <u>184</u>, 185 (1980);
   (b) B. R. Karas and A. B. Ellis, <u>J. Am. Chem. Soc.</u>, <u>102</u>, 968 (1980).
- B. R. Karas, H. H. Streckert, R. Schreiner, and A. B. Ellis, <u>J. Am. Chem.</u>
   <u>Soc.</u>, <u>103</u>, 1648 (1981).
- 4. H. H. Streckert, J. Tong, and A. B. Ellis, submitted for publication.
- N. I. Vitrikhouskii and I. B. Mizetskaya, in "Growth of Crystals," Vol. 3,
   p. 247, Consultants Bureau, New York (1962).
- 6. G. Oelgart, R. Stegmann, and L. John, Phys. Stat. Sol. A., 59, 27 (1980).
- 7. (a) F. L. Pedrotti and D. C. Reynolds, Phys. Rev., 127, 1584 (1962);
  (b) E. T. Handelman and W. Kaiser, J. Appl. Phys. 35, 3519 (1964).
- 8. (a) R. H. Bube, <u>J. Appl. Phys.</u>, <u>35</u>, 586 (1964); (b) Y. S. Park and D. C. Reynolds, <u>Phys. Rev.</u> <u>132</u>, 2450 (1963); (c) R.H. Bube, <u>Phys. Rev.</u>, <u>98</u>, 431 (1955).
- 9. D. Dutton, Phys. Rev., 112, 785 (1958).
- 10. R. B. Parsons, W. Wardzynski, and A.D. Yoffe, Proc. Roy. Soc. A., 262, 120 (1961).
- 11. R. N. Noufi, P. A. Kohl, and A. J. Bard, This Journal, 125, 375 (1978).
- R. E. Halsted in "Physics and Chemistry of II-VI Compounds," M. Aven and
   J. S. Prener, Editors, Chapter 8, North-Holland Publishing Co., Amsterdam (1967).
- 13. Z. Harzion, N. Croitoru, and S. Gottesfeld, This Journal, 128, 551 (1981).
- 14. W. Lehmann, Solid-State Electronics, 9, 1107 (1966).
- M. S. Wrighton, D. S. Ginley, and D. L. Morse, J. Phys. Chem., 78, 2229 (1974).
- 16. H. Gerischer, J. Electroanal. Chem. Interfacial Electrochem., 58, 263 (1975).
- 17. B. R. Karas, D. J. Morano, D. K. Bilich, and A. B. Ellis, <u>This Journal</u>, <u>127</u>, 1144 (1980).
- 18. A. B. Ellis, B. R. Karas, and H. H. Streckert, <u>Faraday Discuss. Chem. Soc.</u>, 70, 165 (1980).

### References cont.

- 19. D. Cahen, B. Vainas, and J. M. Vandenberg, This Journal, 128, 1484 (1981) and references therein.
- (a) K. H. Beckmann and R. Memming, <u>This Journal</u>, <u>116</u>, 368 (1969);
  (b) R. Memming, <u>ibid</u>, <u>116</u>, 785 (1969);
  (c) B. Pettinger, H.-R. Schöppel, and H. Gerischer, Ber. Bunsenges Phys. Chem. 80, 849 (1976).
- 21. (a) A. B. Ellis, S. W. Kaiser, J. M. Bolts, and M. S. Wrighton, <u>J. Am. Chem.</u>
  Soc., <u>99</u>, 2839 (1977); (b) T. Watanabe, A. Fujishima, and K. Honda, <u>Chem.</u>
  Lett., 897 (1974).
- 22. H. H. Streckert, B. R. Karas, D. J. Morano, and A. B. Ellis, <u>J. Phys. Chem.</u> 84, 3232 (1980).
- (a) D. M. Kolb and H. Gerischer, <u>Electrochim. Acta</u> 18, 987 (1973);
  (b) H. Masuda, A. Fujishima, and K. Honda, <u>Chem. Lett.</u>, 1153 (1980).
- 24. (a) A. J. Bard, C. P. Keszthelyi, H. Tachikawa, and N. E. Tokel in "Chemiluminescence and Bioluminescence," M. J. Cormier, D. M. Hercules and J. Lee, Eds. pp. 193-208, Plenum Press, New York (1973); (b) L. R. Faulkner and A. J. Bard, <u>Electroanal. Chem.</u>, <u>10</u>, 1 (1977); (c) C. P. Keszthelyi, N. E. Tokel-Takvoryan, and A. J. Bard, <u>Anal. Chem.</u> <u>47</u>, 249 (1975).
- 25. N. S. Gill and F. B. Taylor, Inorg. Syn. 9, 136 (1967).

Table I. Temperature Dependence of Photoluminescence Spectra

	λ παχ, <sup>n</sup>	λ max,nm(eV) <sup>b</sup>		
CdS <sub>X</sub> Se <sub>1-X</sub> Sample, X	295 K	77 K	Spectral Shift, eV	Intensity Enhancement
1.00 (CdS)	508(2,44)	488(2.54) 502 sh (2.47) 510 sh (2.43) 592 (2.09)	0,10	50
0.74	563(2.20)	545 (2.27) 555 sh(2.23)	0.07	200
0.49	615(2,02)	591 (2.10) 604 sh(2.05)	0°08	200
0.11	695(1,78)	665 (1.86) 680 sh (1.82) 850(1.46)e	0.08	100
0.00 (CdSe)	718(1,73)	681 (1.82) 698 sh (1.78) 830 (1.49)e	60*0	20

# Table I footnotes

- a. Properties of uncorrected PL spectra of  $CdS_XSe_{1-X}$  samples excited with 457.9—nm excitation. Spectra were recorded at 295, 77, and 295 K again to demonstrate reproducibility.
- b. Emission band maxima at 295 and 77 K. At 77 K the highest-energy band generally displayed a shoulder whose position is indicated by "sh". Other bands were occasionally seen and their positions indicated as well. Spectral resolution was ~ 2.0 nm.
- c. Spectral shift between 295 and 77 K for the highest-energy emission band maxima.
- d. Factor by which the emitted intensity of the highest-energy band increased on cooling from 295 to 77 K. This factor has been corrected for relative detector sensitivity (see Experimental).
- e. Recorded using the McPherson-based spectrometer (see Experimental).

Table II. Measurements of Efficiency in Luminescent CdS  $_{
m X}^{
m Se}$ 1- $_{
m X}^{
m -based}$  PECs  $^{
m a}$ 

φ <sub>xcalc.</sub> c φ <sub>xmeas.</sub> d φ <sub>x at ηmax</sub> e 0.41 0.37 0.30 0.93 0.91 0.63	c. $\phi_{\text{x}}$ d $\phi_{\text{x}}$ at $\eta_{\text{max}}$ 0.30
	6
	E <sub>v</sub> at n <sub>max</sub> , v 0.42

### Table II footnotes

- a. PEC characterization parameters extracted from the iLV curves of Figure 5. The indicated crystal served as the photoanode in a PEC employing  $N_2$  purged  $1\,M\,OH^-/1\,M\,S^{2-}/0.1\,M\,S$  polysulfide electrolyte; a Pt foil counterelectrode and SCE completed the PEC. Electrodes were irradiated with  $\sim 1.0$  to  $1.2\,$  mW of 457.9 -nm excitation; their exposed surface areas were  $\sim 0.25\,$  cm $^2$ .
- b. Ratio of open-circuit to in-circuit emission intensity with the in-circuit value taken at 0.30 V vs SCE. Emission intensity was monitored at the emission band maximum.
- c. Photocurrent quantum efficiency at 0.30 V vs. SCE calculated from the  $\phi_{r_0}/\phi_{r_i}$  ratio in the preceding column and eq. (2) in the text.
- d. Measured photocurrent quantum efficiency at 0.30 V vs. SCE determined with a radiometer as described in the Experimental. Measured values are uncorrected for reflection and electrolyte absorption.
- e. Measured photocurrent quantum efficiency at the potential where maximum optical to electrical energy conversion efficiency obtains.
- f. Output voltage at the maximum energy conversion point. The redox potential of the polysulfide electrolyte was - 0.78 V vs. SCE.
- g. (Maximum electrical power out divided by input optical power) x 100.
- the others and would, therefore, require a larger correction for electrolyte absorption.

Table III. Estimates of Electroluminescence Efficiency

0.00 (CdSe)	0.11	0.49	0.74	1.00 (CdS)	CdS <sub>X</sub> Se <sub>1-X</sub> F Electrode,X V
-1.50	-1.50	-1.50	-1.50	-1.50	Potential,
-1.00	-1.00	-1.00	-1.00	-1.05	V vs. SCE <sup>b</sup>
5.6 (34)	1.2 (7.1)	0.61 (3.2)	0.23 (1.1)	0.11 (0.49)	Total emitted light, µJ (10 <sup>12</sup> einsteins) <sup>C</sup>
0.056 (0.34)	0.036 (0.21)	0.0014 (0.0073)	0.0029 (0.014)	0.00063 (0.0027)	
7.0 (7.2)	5.7 (5.9)	9.7 (10.1)	6.8 (7.1)	3.4 (3.5)	10 <sup>3</sup> total Coul (10 <sup>8</sup> mole e <sup>-</sup> ) <sup>d</sup>
0.53 (0.55)	0.40 (0.42)	0.080 (0.083)	0.61 (0.63)	0.049 (0.050)	
4.7 0.62	1.2 0.51	0.32	0.15 0.023	0.14 0.054	10 <sup>4</sup> $\overline{\phi}_{EL}$

### Table III footnotes

- a. A single-crystal, n-type  $\mathrm{CdS}_{X}\mathrm{Se}_{1-X}$  sample was used as the working electrode in a one-compartment EL experiment conducted in a N<sub>2</sub>-purged 5 M NaOH/0.1 M K<sub>2</sub>S<sub>2</sub>O<sub>8</sub> electrolyte; a Pt foil counterelectrode and SCE completed the cell. The  $\mathrm{CdS}_{X}\mathrm{Se}_{1-X}$  exposed surface areas were  $\sim 0.25~\mathrm{cm}^2$ .
- b. EL experiments were conducted by pulsing the  $CdS_XSe_{1-X}$  electrode between 0.00 V (11 s) and the indicated potential (1 s).
- c. Total emitted light collected per pulse with a radiometer operated in its integrating mode. The radiometer was placed ~1 cm away from the emitting electrode. If all of the indicated energy were converted to photons at the emission band maxima (508, 563, 615, 695 and 718 nm for X = 1.00, 0.74, 0.49, 0.11, and 0.00, respectively), the quantity of einsteins given in parentheses would be obtained.
- d. Total coulombs collected per pulse with a digital coulometer. The value in parentheses is the corresponding quantity of electrons in the external circuit, obtained by dividing the coulombs by Faraday's constant.
- e. Integrated EL efficiency. This value is derived by dividing the einsteins by the moles of electrons from the preceding columns. These efficiencies are lower limits, as described in the text. Table entries for each potential are representative values. At least five pulse sequences were used at each potential, yielding  $\overline{\phi}_{\rm EL}$  values within 10% of one another.

## Figure Captions

- Figure 1. Absorbance spectra of single-crystal CdS Se  $_{\rm X}$  samples. Thicknesses are 0.10, 0.14, 0.12, 0.13, and 0.13 cm for X = 1.00, 0.74, 0.49, 0.11, and 0.00, respectively. The samples were polished with 1 - $\mu$ m alumina.
- Figure 2. (a) Uncorrected PL spectra of CdS<sub>X</sub>Se<sub>1-X</sub> samples irradiated in 5 M OH electrolyte while they were held at -1.50 V vs. SCE. The ~ 0.25-cm<sup>2</sup> samples were excited with ~ 1.0 mW of 457.9-nm light (excitation spike shown at 1/100 the scale of the PL spectrum). Emission intensities are not directly comparable because of differences in geometry. (b) Uncorrected EL spectra of the samples in (a) obtained without changing their geometry. The electrolyte in the EL experiments was 5 M OH /0.1 MS<sub>2</sub> 0<sub>8</sub><sup>2-</sup>. Electrodes were continuously pulsed between 0.00 V (11s) and 1.50 V vs. SCE (1s) while the emission spectrometer was scanned at 12 nm/min. For both PL and EL a spectral resolution of 2.0 nm was employed.
- Figure 3. Uncorrected PL (a) and EL (b) spectra of  $CdS_XSe_{1-X}$  taken as Jascribed in the Figure 2 caption.
- Figure 4. Plot of the emission band maxima (nm) from Figures 2a and 3a vs. composition, X, of the  $CdS_{\chi}Se_{1-\chi}$  samples.
- Figure 5. Photocurrent (solid lines, left-hand scales) and emission intensity (dashed lines, right-hand scales) monitored at emission band maxima for single-crystal CdS<sub>X</sub>Se<sub>1-X</sub> electrodes in 1 M OH<sup>-</sup>/1 M S<sup>2-</sup>/0.1 M S electrolyte as a function of potential. The electrodes (~0.25-cm<sup>2</sup> exposed area) were excited with ~1.0-1.2 mW of 457.9-nm light from a beam-expanded Ar ion laser. These iLV curves were swept at 10 mV/s. The electrolyte redox potential was -0.78 V vs. SCE.

- Figure 6. Current-voltage curves for CdS<sub>0.49</sub>Se<sub>0.51</sub>, CdS<sub>0.11</sub>Se<sub>0.89</sub>, and CdS<sub>0.74</sub>Se<sub>0.26</sub>, labelled 1, 2, and 3, respectively. Dashed lines represent data obtained in 5 M NaOH electrolyte; solid lines are plots obtained in 5 M NaOH, 0.1 MK<sub>2</sub>S<sub>2</sub>O<sub>8</sub> electrolyte.

  All curves were swept at 20 mV/s. Electrode surface areas exposed to the electrolytes were ~0.25 cm<sup>2</sup>.
- Figure 7. Uncorrected PL (solid curve) and EL (vertical lines) spectra of  $CdS_{0.49}Se_{0.51}$  obtained in the same sample geometry. The PL spectrum was taken with the electrode immersed in 5 MNaOH electrolyte at 1.50 V vs. SCE. The 514.5-nm line of an Ar ion laser whose beam was expanded and masked to fill the  $\sim 0.25$ -cm<sup>2</sup> electrode surface was used for excitation; the excitation spike is shown in the inset. The EL spectrum was taken in 5 M NaOH/0.1 M K<sub>2</sub>S<sub>2</sub>O<sub>8</sub> electrolyte. The electrode was continuously pulsed between 0.00 V (11 s) and -1.50 V vs. SCE (1 s) while the emission spectrum was scanned at 6 nm/min. For both PL and EL a spectral resolution of 0.3 nm was employed. The EL spectrum has been scaled down to match the PL intensity at λ<sub>max</sub>  $\sim 615$  nm.
- Figure 8. Relative emission intensity vs. potential for CdS<sub>X</sub>Se<sub>1-X</sub> electrodes in 5 M NaOH electrolyte. Curves 1,2, and 3 correspond to CdSe, CdS and CdS<sub>0.49</sub>Se<sub>0.51</sub>, respectively. The samples were all irradiated with ~0.1-3 mW of 457.9 nm light; the laser beam was expanded and masked to fill the ~0.25-cm<sup>2</sup> sample surface areas. Because the geometries of the three samples were not identical relative to the detection optics, the intensities between curves are not comparable. The potential was swept in all cases at 20 mV/s beginning at -0.90 V vs. SCE.

















